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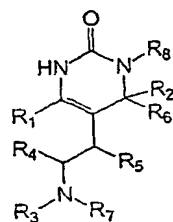
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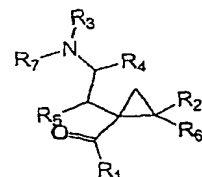
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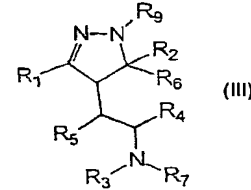
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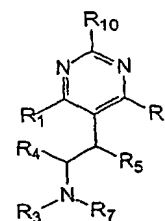
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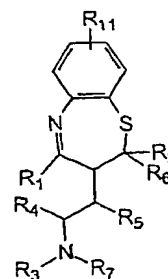
(II)



(III)



(IV)



(V)

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(57) Abstract: The present invention provides a combinatorial approach to a library of novel compounds having four diversity points. The compounds provide for the mapping of urotesin II and somatostatin 5 receptors by differential binding of said receptors. Accordingly, the invention relates in a first aspect to novel compounds of the general formula I to V or salts thereof. The present invention further relates to a method of treating diseases for which modulation of the urotesin II receptor produces a physiologically beneficial response in said disease, such as those associated with CNS function and cardiovascular diseases. The present invention further relates to pharmaceutical compositions comprising these agents for the treatment of these diseases adapted to modulate the urotesin II receptor.